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CDC's Universal Data Collection System

Report on the Universal Data Collection System (UDC)

Includes data collected from
May 1998 through October 1999



U. S. DEPARTMENT OF HEALTH AND HUMAN SERVICES
Centers for Disease Control and Prevention
Atlanta, Georgia 30333



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Single copies of the *Report on the Universal Data Collection System* are available free from HANDI, the information service of the National Hemophilia Foundation by calling (800) 42-HANDI. Confidential information, referrals, and educational material on hemophilia and other bleeding disorders is also available through HANDI. The *Report on the Universal Data Collection System* is accessible via internet at <http://www.cdc.gov/ncidod/dastlr/Hematology/HDBarchive.htm>.

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Commentary

The two most common congenital bleeding disorders are von Willebrand disease (vWD) and hemophilia. vWD is caused by defective synthesis or function of a protein, called von Willebrand factor, which is necessary for normal blood clotting. vWD occurs with equal frequency in men and women. Although the prevalence of this disease is not precisely known, it is estimated that between one and two percent of the population are affected. There are different types and severity of vWD. Symptoms include heavy or prolonged menstrual bleeding, easy bruising, frequent or prolonged nosebleeds, and prolonged bleeding following surgery, dental work, childbirth, or injury.

Hemophilia is caused by a defect in the gene located on the X chromosome that contains the genetic code for one of the clotting factor proteins necessary for normal blood clotting. A deficiency of factor VIII is referred to as hemophilia A or “classic” hemophilia. In contrast, a deficiency of factor IX characterizes hemophilia B, also known as Christmas disease. The defect usually occurs on one of the two female X chromosomes and results in a carrier state. When males have the defect on their only X chromosome, they are affected with the disease. Thus, almost all of the approximately 17,000 persons with hemophilia in the United States are male.

People with severe hemophilia can experience serious bleeding into tissues, muscles, joints, and internal organs, often without any obvious trauma. Repeated bleeding into joints without adequate treatment results in crippling chronic joint disease, one of the severe complications of bleeding disorders. In the mid-1970s, treatment for hemophilia was improved through the use of clotting factor concentrates, products made from the plasma of donated blood. However, because blood donations from thousands of donors are pooled together to make these products, many persons with bleeding disorders were

infected with hepatitis B and C viruses and with human immunodeficiency virus (HIV), the virus that causes AIDS, before the risk of disease transmission in blood products was recognized and prevention measures taken.

In 1975, Congress initiated federal funding to specialized hemophilia treatment centers (HTCs) to provide comprehensive care to persons with bleeding disorders. Since 1986, CDC has been involved with the hemophilia community through the HTC system, primarily through risk-reduction efforts aimed at preventing secondary infection of family members with HIV.

In 1991, CDC received a request from the National Hemophilia Foundation to expand their collaborative activities within the bleeding disorders community. Meetings with patients and hemophilia care providers were held during 1992 to determine the areas of highest priority. Based on recommendations from these constituents, a Congressional mandate was issued to CDC, with the goal of reducing the human suffering and financial burden of bleeding disorders by focusing national emphasis on prevention and early intervention. The issues of greatest concern identified by the bleeding disorders community were: 1) the safety of the blood supply from infectious diseases; and 2) the prevention of joint disease.

In response, CDC developed the Universal Data Collection System (UDC). The purpose of UDC is two-fold: 1) to establish a sensitive blood safety monitoring system among persons with bleeding disorders; and 2) to collect a uniform set of clinical outcomes information that could be used to monitor the occurrence of and potential risk factors for infectious diseases and joint complications.

Persons with bleeding disorders are enrolled in UDC by care providers in each of the nation's 134 federally funded HTCs. As part of the project, a uniform set of clinical data and a plasma specimens are collected by HTC

staff each year during the participant's annual comprehensive clinic visit. A portion of the plasma specimen is used to perform free screening tests for hepatitis A, B, and C viruses and for HIV. The remainder of the specimen is stored for use as needed in future blood safety investigations.

Enrollment in UDC began in May 1998. Information about eligibility requirements, enrollment procedures, and data collection can be found in the *Technical Notes* of this report. Participating HTC's are listed by region in the *Acknowledgements*. A regional map is included at the end of this report.

The purpose of this surveillance report is to disseminate the information being collected by this project to public health workers, health educators and planners, other care providers, and patients in the bleeding disorders community. The report contains information about the demographic characteristics of the participants, their blood and factor product use, and the occurrence and treatment of joint and infectious diseases. We hope that this information will prove useful to those involved in efforts to reduce or prevent the complications of these conditions.

The proper interpretation and appropriate use of surveillance data require an understanding of how the data are collected, reported, and analyzed. Therefore, readers of this report are encouraged to review the *Technical Notes*, beginning on page 18.

Suggested Reading:

CDC. Prevention of hepatitis A through active or passive immunization. Recommendations of the Advisory Committee on Immunization Practices (ACIP). *MMWR* 1996;45(No. RR-15):1-30.

CDC. Transmission of hepatitis C virus infection associated with home infusion therapy for hemophilia. *MMWR* 1997;46:597-599.

CDC. Occurrence of hemophilia in the United States. *American Journal of Hematology* 1998; 59:288-294.

Hill H, Stein S. Viral infections among patients with hemophilia in the state of Georgia. *American Journal of Hematology* 1998;59:36-41.

The following publications are available from HANDI (800-42-HANDI):

- *What You Should Know about Bleeding Disorders* (1997)

- *Comprehensive Care for People with Hemophilia* by Shelby Dietrich, MD (1991)

- *Understanding Hepatitis* by Leonard Seeff, MD (1997)

- *HIV Disease in People with Hemophilia: Your Questions Answered* by Glenn Pierce, MD, PhD (1991)

- *Bleeding Disorders and AIDS: The Facts* (1997)

- Information packet on von Willebrand disease.

Table 1. Enrollment in UDC, May 1998 – October 1999

<u>Month</u>	<u>Number Enrolled</u>	<u>Number Refused</u>	<u>Refusal Rate (%)</u>
May – Oct, 1998	988	85	7.9
November	224	25	10.0
December	219	26	10.6
January, 1999	318	25	7.3
February	359	38	9.6
March	368	49	11.8
April	363	40	9.9
May	359	43	10.7
June	415	40	8.8
July	307	42	12.0
August	355	64	15.3
September	297	54	15.4
October	236	43	15.4
Total	4808	574	10.7

Table 2. Regional* enrollment activity, May 1998 – October 1999

<u>Region</u>	<u>Number Approached</u>	<u>Refusal rate (%)</u>
I	216	11.6
II	776	12.6
III	805	16.4
IV-N	480	6.3
IV-S	308	10.7
V-E	289	11.8
V-W	633	7.6
VI	628	13.1
VII	206	11.7
VIII	327	3.7
IX	634	7.9
X	14	0

*See map (page 24) for regional designations.

Table 3. Demographic characteristics of persons* enrolled in UDC

Characteristic	Hemophilia				vWD	
	A (n = 3248)		B (n = 761)		(n = 723)	
	Number	Percent	Number	Percent	Number	Percent
Age Group (years)						
2 – 10	918	28.4	192	25.3	199	27.6
11 – 20	1007	31.1	199	26.3	262	36.3
21 – 40	798	24.6	229	30.2	131	18.1
41 – 60	443	13.7	108	14.2	93	12.9
61+	72	2.2	30	4.0	37	5.1
Race / Ethnicity						
White	2236	68.9	556	73.1	529	73.2
African American	391	12.0	92	12.1	28	3.9
Hispanic	416	12.8	81	10.6	69	9.5
Asian / Pacific Islander	78	2.4	8	1.1	21	2.9
Native American	27	0.8	6	0.8	7	1.0
Other	99	3.0	18	2.4	69	9.5
Sex						
Male	3189	98.2	743	97.6	342	47.3
Female	59	1.8	18	2.4	381	52.7

*Twenty-four persons were reported to have both hemophilia and vWD, and 98 persons had a bleeding disorder other than hemophilia or vWD.

Table 4. Disease severity of persons enrolled in UDC

	Hemophilia						vWD			
	Mild		Moderate		Severe		Type 1 & 2		Type 3	
	N	%	N	%	N	%	N	%	N	%
Participants*	821	20.5	919	23.0	2261	56.5	580	88.5	75	11.5

*Numbers do not equal total number of persons because of missing data.

Table 5. Bleeding episodes* among persons enrolled in UDC by disease severity and prophylaxis status

No prophylaxis

Bleeding site	Hemophilia			vWD	
	Mild n = 817	Moderate n = 825	Severe n = 1687	Type 1 & 2 n = 560	Type 3 n = 74
Joint	0.6 (2.3)	3.8 (7.8)	9.3 (12.1)	0.2 (1.6)	3.7 (8.3)
Muscle	0.3 (1.0)	1.1 (2.4)	2.6 (5.8)	0.1 (0.8)	0.5 (1.3)
Other	0.7 (2.9)	1.4 (4.4)	2.0 (4.7)	3.1 (8.4)	3.9 (6.5)
All sites					
Mean	1.7 (4.1)	6.2 (10.3)	13.9 (16.1)	3.4 (8.6)	8.0 (10.5)
Median	0	3	10	0	4

With prophylaxis

Bleeding site	Hemophilia		
	Mild N = 3	Moderate n = 49	Severe n = 315
Joint	0.5 (1.0)	2.7 (4.0)	3.2 (5.8)
Muscle	0 (—)	1.1 (5.4)	0.8 (2.2)
Other	1.2 (1.5)	1.2 (3.3)	1.4 (6.4)
All sites			
Mean	1.8 (1.3)	5.1 (9.3)	5.4 (9.5)
Median	2	3	2

*Values are mean (\pm SD) number of bleeding episodes experienced during the 6-month period preceding the UDC visit.

Table 6. Blood and factor products used* by persons enrolled in UDC

Treatment product	Hemophilia A		Hemophilia B		vWD	
	Number	Percent	Number	Percent	Number	Percent
Recombinant factor	1991	61.3	406	53.4	4	0.6
Monoclonal factor VIII	664	20.4	1	0.1	2	0.3
Other human factor [†]	156	4.8	1	0.1	148	21.2
Porcine factor VIII	6	0.2	0	—	0	—
Purified factor IX	2	0.1	277	36.4	0	—
Prothrombin complex	59	1.8	32	4.2	0	—
Activated prothrombin complex	156	4.8	11	1.4	0	—
Cryoprecipitate or FFP	12	0.4	5	0.7	14	2.0
Desmopressin	191	5.9	2	0.3	263	37.6
None used	294	9.1	122	16.0	259	37.1

*Any use of the product(s) during the 12-month period preceding UDC enrollment.

NOTE: Individuals may have used more than one type of treatment product.

Table 7. Infectious disease complications among persons enrolled in UDC

Infectious Disease Complications	Hemophilia		vWD	
	Number	% of Total	Number	% of Total
Risk factors for liver disease				
Past/present hepatitis B virus infection	808	20.2	25	3.5
Past/present hepatitis C virus infection	1883	47.0	59	8.2
History of alcohol abuse	147	3.7	3	0.4
Other	41	1.0	8	1.1
None	2033	50.7	650	89.9
Signs or symptoms of liver disease (During the last year)				
Jaundice	27	0.7	1	0.1
Ascites	25	0.6	1	0.1
Varices	17	0.4	0	—
Other	45	1.1	1	0.1
None	3920	97.8	720	99.6
Laboratory markers of liver disease				
Chronically elevated ALT/AST levels	721	18.0	15	2.1
Elevated prothrombin time in the last year	101	2.5	10	1.4
Therapy for chronic viral hepatitis				
Any therapy	167	4.2	5	0.7
Successful therapy	30	18.0*	1	20.0*
Intravenous access devices (IVAD)				
Used an IVAD in the last year	496	12.4	24	3.3
IVAD infection in the last year	66	13.3**	3	12.5**

*Percent of persons who received any therapy for chronic viral hepatitis.

**Percent of persons who used an IVAD in the last year.

Figure 1. Prevalence of hepatitis A virus exposure and reported vaccination status among persons with hemophilia

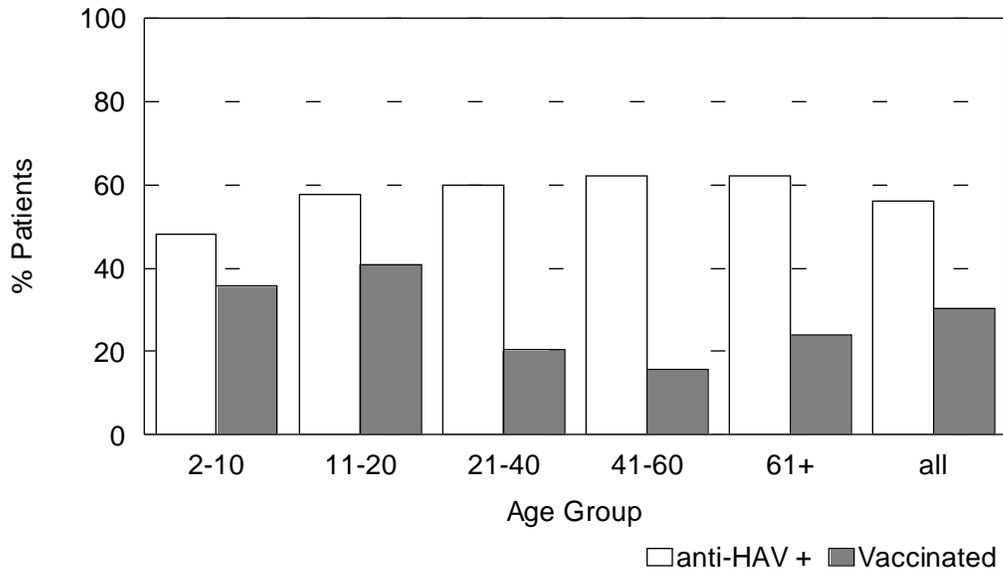
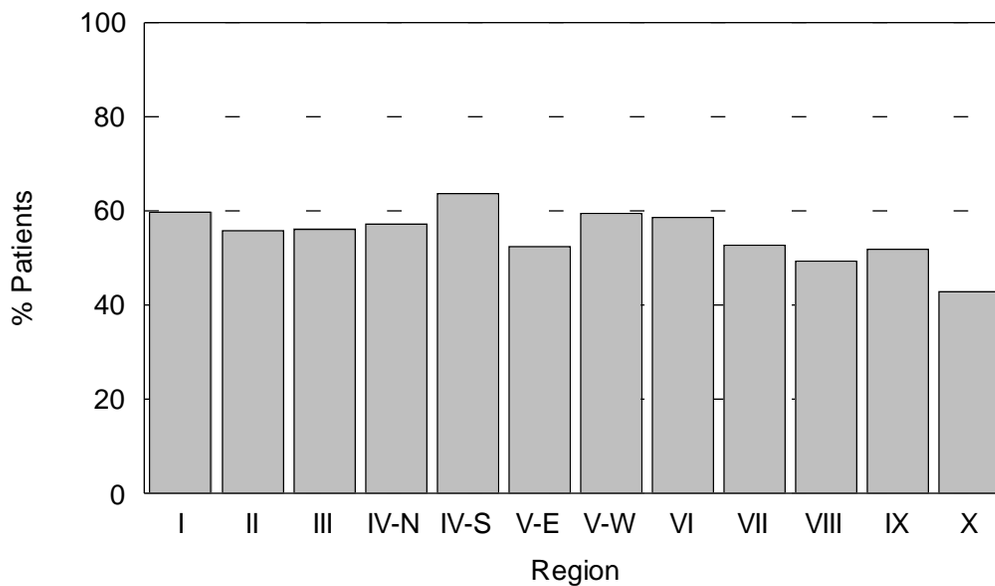


Figure 2. Regional* distribution of natural and acquired immunity to hepatitis A virus among persons with hemophilia



*See map (page 24) for regional designations.

Figure 3. Prevalence of hepatitis A virus exposure and reported vaccination status among persons with vWD

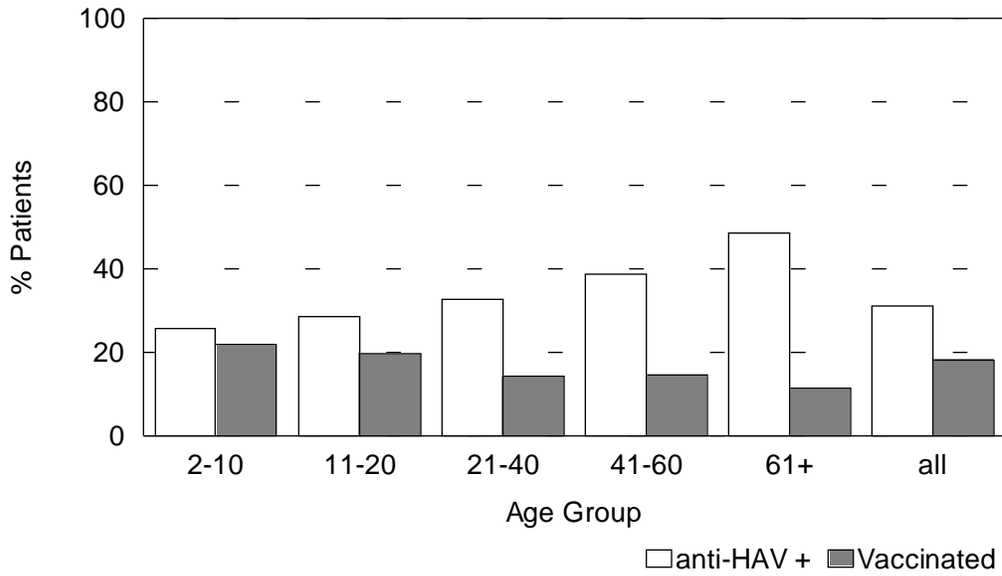


Figure 4. Prevalence of hepatitis B virus exposure and reported vaccination status among persons with hemophilia

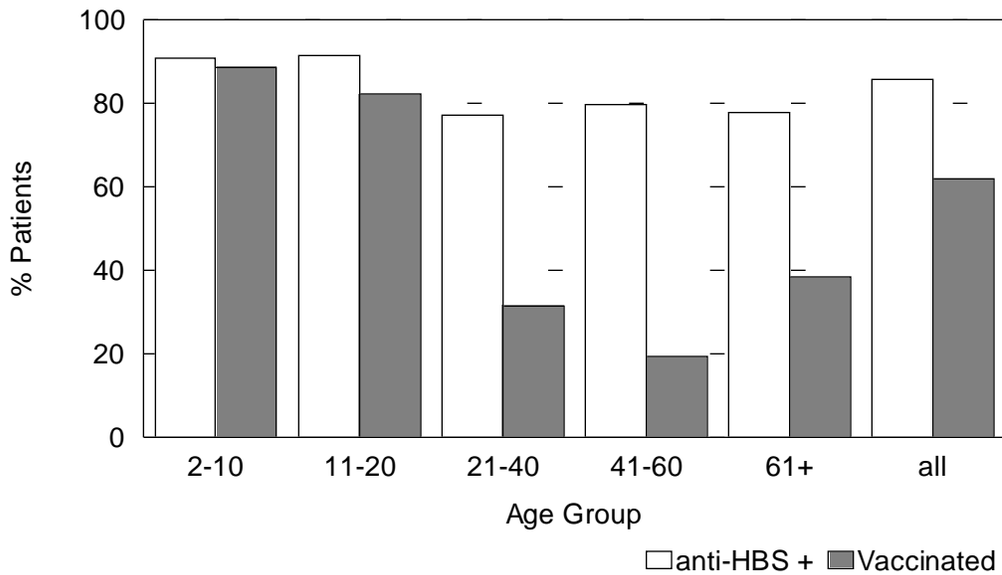
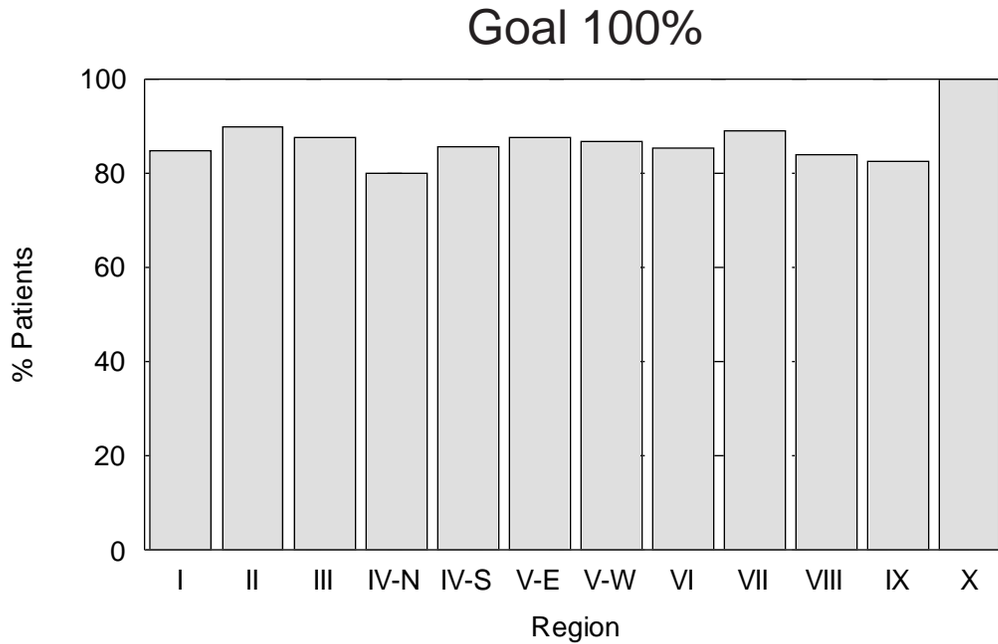


Figure 5. Regional* distribution of natural and acquired immunity to hepatitis B virus among persons with hemophilia



*See map (page 24) for regional designations.

Figure 6. Prevalence of hepatitis B virus exposure and reported vaccination status among persons with vWD

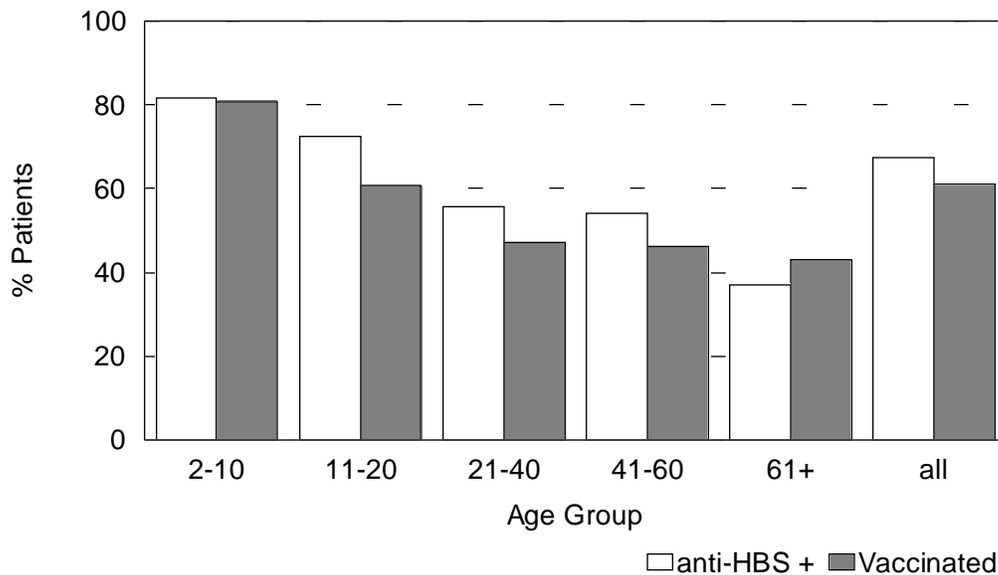


Figure 7. Prevalence of hepatitis C virus infection among persons with bleeding disorders

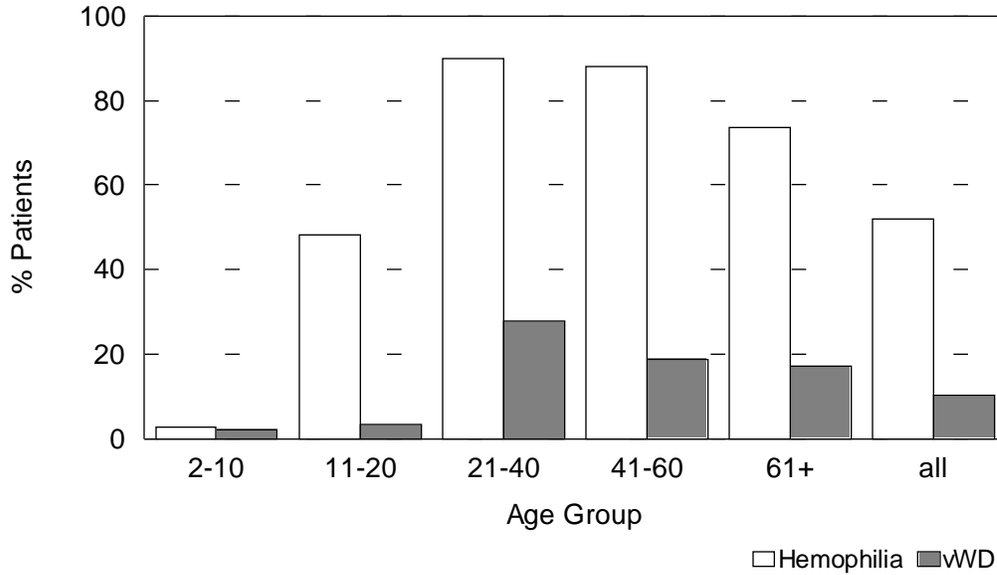


Figure 8. Prevalence of human immunodeficiency virus (HIV) infection among persons with bleeding disorders

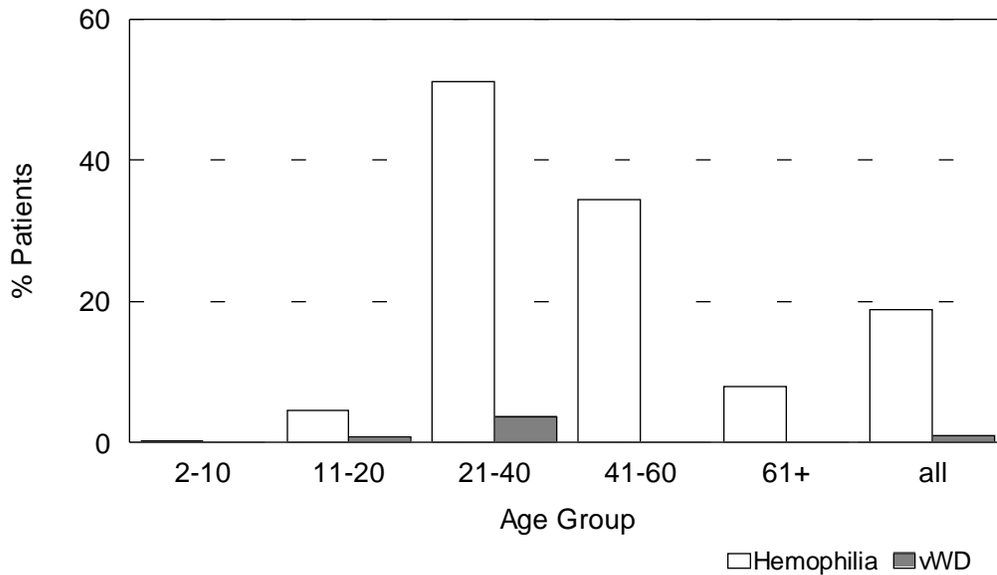


Table 8. Treatment type for persons with hemophilia enrolled in UDC

Severity	Total number	Episodic care No. (%)	Number on intermittent prophylaxis	Number on continuous* prophylaxis
Mild	821	808 (98.4)	9	4
Moderate	919	784 (85.3)	41	94
Severe	2261	1503 (66.5)	186	572

*Prophylaxis is considered continuous when administered for at least 46 weeks per year.

Table 9. Prevalence of current inhibitors by titer* among persons with hemophilia enrolled in UDC

Severity	Hemophilia A			Hemophilia B		
	Number	Low titer	High titer	Number	Low titer	High titer
Mild	650	2 (0.1)	1 (0.03)	171	0	0
Moderate	659	11 (1.7)	9 (1.4)	260	0	1 (0.4)
Severe	1932	74 (3.8)	116 (6.0)	329	4 (1.2)	15 (4.6)

*Low titer is defined as an inhibitor level of 0.5 – 5 Bethesda units (BU).
 High titer is defined as an inhibitor level of >5 BU.
 Numbers in parentheses are percents.

Table 10. Intra-cranial hemorrhage (ICH)* among persons with hemophilia enrolled in UDC

Severity	Hemophilia A		Hemophilia B	
	Total	No. with ICH (%)	Total	No. with ICH (%)
Mild	650	1 (0.2)	171	0
Moderate	659	6 (0.9)	260	1 (0.4)
Severe	1932	21 (1.1)	329	4 (1.2)
Causes of ICH			Number (%)	
Trauma			15 (53.6)	
Thrombocytopenia			1 (3.6)	
Other			12 (42.9)	

*Diagnosed by a physician during the year prior to the UDC visit.

Table 11. Joint complications among persons enrolled in UDC

	Hemophilia						vWD			
	Mild		Moderate		Severe		Type 1 & 2		Type 3	
	N	%	N	%	N	%	N	%	N	%
Target joint*	73	8.9	281	30.6	1129	49.9	10	1.8	17	22.7
Invasive procedure	33	4.0	63	6.9	277	12.2	9	1.6	7	9.3
Joint infection	6	0.7	14	1.5	39	1.7	3	0.5	0	—
Used cane	114	13.9	230	25.0	711	31.4	35	6.2	16	21.3
Used wheelchair	14	1.7	60	6.5	236	10.4	15	2.7	4	5.3
Any activity restriction	119	14.5	282	30.7	957	42.3	40	7.1	21	28.0

*Please see Technical Notes (page 18) for the definition of a target joint

Table 12. Joint limitations among persons enrolled in UDC

	Hemophilia			vWD	
	Mild	Moderate	Severe	Type 1 & 2	Type 3
Number of patients	777	841	1992	531	70
Mean indicator* value	53.7	96.9	159.4	28.2	67.0
Standard deviation	99.4	175.7	220.8	67.0	110.3

*Indicator is the total number of degrees of range of motion less than normal for five joints. The joint motions measured and normal values used (in parentheses) are: hip extension (30); hip flexion (120); knee flexion (135); knee extension (0); shoulder flexion (180); elbow flexion (150); elbow extension (0); elbow pronation and supination (80); ankle dorsiflexion (20); ankle plantar flexion (50). Any hyperextension of the knee or elbow is not included in the calculation. In UDC, limitations in knee and elbow extension are recorded as negative numbers. Patients with missing measures for any of the joints are excluded from the analyses. As an example, patients with mild hemophilia have on average 53.7 degrees less than normal range of motion across ten joints.

Technical Notes

Eligibility Requirements

To participate in UDC, patients must receive care in a federally funded HTC and meet at least one of the following criteria: 1) age 2 years or older with a bleeding disorder due to congenital deficiency or acquired inhibitors in which any of the coagulation proteins is missing, reduced, or defective and has a functional level of less than 50 percent; or 2) age 2 years or older with a diagnosis by a physician of von Willebrand disease. Individuals specifically excluded from participation in UDC include persons with any of the following: 1) an exclusive diagnosis of a platelet disorder; 2) thrombophilia; or 3) coagulation protein deficiencies due to liver failure.

Data collection

UDC data are collected during the participant's "annual visit," which ideally should occur once each calendar year (January-December), with the interval between visits as close as possible to 12 months. Data are collected according to guidelines and definitions detailed in surveillance manuals provided to HTC staff by CDC. Informed consent for participation is obtained each year. Demographic information and reasons for refusal are obtained using a Patient Refusal Form for all eligible persons who decline participation. To protect patient confidentiality, all data sent to CDC do not contain personally identifying information, but rather use a unique 12-digit code that is generated by a computer software program supplied to HTCs by CDC.

Eligible participants are registered into UDC through a Registration Form completed by HTC staff; this form includes patient demographic, diagnostic, and historical information. Month and year of birth are used to calculate age on the last day of the current year. Information on race and ethnicity is obtained from

clinic records and may have been based either on self-report or on observations made by care providers.

During the annual visit, clinical information is recorded on a standardized data collection form (Annual Visit Form). In addition to information about education, employment status, and health insurance, data are also collected about treatment type (episodic vs. prophylactic), presence and treatment of inhibitors, the number of bleeding episodes experienced (based on infusion logs or patient recall), type and brand name of all factor concentrates or other treatment products used, and whether or not clotting factor is infused at home.

Information regarding infectious diseases is also collected including risk factors and clinical signs, symptoms, and laboratory markers of liver disease. Data are also recorded about any therapy for chronic hepatitis, the status of vaccination for hepatitis A and B viruses, and, among patients with an intravenous access device, the occurrence of a device-associated infection. Persons ≥ 16 years of age who are HIV-infected are asked several questions concerning risk-reduction activities including partner testing and condom use.

Data are also collected on joint disease, including the use of walking aids, the occurrence of joint infections, and measures of impact of joint disease on daily activities. During the visit, range of motion measurements on five joints (hip, knee, shoulder, elbow, and ankle) are taken by a physical therapist or other trained health care provider according to detailed guidelines provided in a reference manual supplied by CDC. All health care providers performing these measurements are trained and certified by regional physical therapists who have themselves received centralized training. In addition, information about whether a particular joint is a "target joint" or whether the participant has

required the use of an orthopedic appliance or has undergone an invasive orthopedic procedure is collected. In UDC, a target joint is defined as a joint in which recurrent bleeding has occurred on four or more occasions during the previous 6 months or one in which 20 life-time bleeding episodes have occurred.

All data collection forms are sent overnight to CDC where they are then key entered into a computer database using double-entry software to minimize data entry errors. Data are then screened for omissions, inconsistencies, and unusual values that possibly represent abstraction or data-entry errors. Error reports are generated and faxed to the HTC, where a designated UDC contact uses available information to resolve discrepancies and complete missing data items.

Laboratory testing

During the annual visit, a blood specimen is obtained from each participant in UDC. The specimen is processed by HTC personnel according to guidelines provided by CDC that are designed to minimize the effects of storage and shipment on subsequent analyses. Samples are shipped overnight to the CDC Serum Bank where they are aliquoted and stored. A portion of the specimen is sent to the Eugene B. Casey Hepatitis Laboratory at Baylor College of Medicine in Houston, Texas. A second portion is sent to the HIV Testing Laboratory at CDC. The remainder of the specimen is stored in the CDC Serum Bank for future blood safety investigations, as needed.

Testing for hepatitis A, B, and C viruses follow algorithms designed to determine with the highest probability the patient's status with regard to exposure to or infection with these viruses. Information provided by HTC staff on a Laboratory Form, including the results of previous local testing and vaccination history, is used by personnel at the testing laboratory to provide a detailed interpretation of the test results.

Testing for HIV follows algorithms designed to determine patient status with regard to infection with HIV-1 and HIV-2. The results of all laboratory testing are reported to the HTC using the CDC unique code which can be matched to the patient only by HTC staff.

Mortality reporting

Deaths occurring among all HTC patients (regardless of whether or not they have been enrolled in UDC) are reported to CDC using a Mortality Form. Data collected include age at death, sex, race/ethnicity, disease type and severity, and whether or not blood products had been used during the year prior to death. Additionally, information about the death, including the date, cause (primary and contributing), and whether or not an autopsy was performed, is also collected.

Tabulation and presentation of data

Data in this report are provisional. The data presented in this report represent the first 18 months of what is planned to be at least a 5-year surveillance project. Future reports will include expanded data tables to cover subsequent surveillance periods and will provide the results of more detailed analyses of available data and findings from special studies.

Acknowledgements

We thank the Regional Coordinators (listed below in italics) of the federal HTC regions for their assistance in the implementation and technical support of UDC. Data for this report were collected by care providers in HTCs at the following institutions:

Region I

Ann Forsberg, M.A., M.P.H.

New England Hemophilia Center
Worcester, MA

Dartmouth-Hitchcock Hemophilia Center
Lebanon, NH

Rhode Island Hospital
Providence, RI

UCONN Hemophilia Treatment Center
Farmington, CT

Vermont Regional Hemophilia Center
Burlington, VT

Boston Children's Hospital
Boston, MA

Region II

Mariam Voutsis, R.N., M.P.A.

The New York Hospital
New York, NY

Puerto Rico Hemophilia Treatment Center
San Juan, PR

UMDNJ-Robert Wood Johnson University
Hospital, New Brunswick, NJ

Nadeene Brunini Comprehensive
Hemophilia Care Center, Newark, NJ

The Mary M. Gooley Hemophilia Center, Inc.
Rochester, NY

SUNY Health Science Center - Adult
Syracuse, NY

SUNY Health Science Center - Pediatric
Syracuse, NY

Hemophilia Center of Western New York –
Adult, Buffalo, NY

Hemophilia Center of Western New York –
Pediatric, Buffalo, NY

Albany Medical College
Albany, NY

UHSB Blood Disorders Center
Johnson City, NY

Mount Sinai Medical Center
New York, NY

Long Island Jewish Medical Center
New Hyde Park, NY

Region III

Sue Cutter, M.S.W., M.P.A.

Children's National Medical Center
Washington, DC

Georgetown University Medical Center
Washington, DC

University of Virginia Hospital
Charlottesville, VA

Medical College of Virginia Hospital
Richmond, VA

Children's Hospital of the King's Daughters
Norfolk, VA

Cardeza Foundation Hemophilia Center
Philadelphia, PA

Christiana Care Health Services
Newark, DE

Hemophilia Center of Central Pennsylvania
Hershey, PA

Hemophilia Center of Western Pennsylvania
Pittsburgh, PA

West Virginia University Medical Center
Morgantown, WV

Charleston Area Medical Center
Charleston, WV

Johns Hopkins University Medical Center
Baltimore, MD

Children's Hospital of Philadelphia Speciality
Center, Voorhees, NJ

Children's Hospital of Philadelphia
Philadelphia, PA

Lehigh Valley Hospital
Allentown, PA

Region IV-N

Richard J. Atwood, M.A., M.P.H.
 Wake Forest University School of Medicine
 Winston-Salem, NC
 Brown Cancer Center
 Louisville, KY
 Children's Hospital of Palmetto –
 Richland Memorial, Columbia, SC
 University of Tennessee – Memphis
 Memphis, TN
 East Tennessee Comprehensive Hemophilia
 Center, Knoxville, TN
 Vanderbilt University Medical Center
 Nashville, TN
 University of North Carolina at Chapel Hill
 Chapel Hill, NC
 Norton Kosair Children's Medical Center
 Louisville, KY

Region IV-S

Crystal D. Watson, B.S.W.
 Miami Comprehensive Hemophilia Center –
 Pediatrics, Miami, FL
 University of Florida
 Gainesville, FL
 Scottish Rite Children's Medical Center
 Atlanta, GA
 Medical College of Georgia - Adult
 Augusta, GA
 University of Mississippi Medical Center
 Jackson, MS
 Miami Comprehensive Hemophilia Center –
 Adult, Miami, FL
 Children's Rehabilitation Services
 Mobile, AL
 Children's Rehabilitation Services
 Birmingham, AL
 Children's Rehabilitation Services
 Opelika, AL
 Children's Rehabilitation Services
 Huntsville, AL
 Emory University Hemophilia Program
 Office
 Atlanta, GA
 University of South Florida – Pediatric
 Tampa, FL

Region V-E

Tamara Wood-Lively, M.H.A., J.D.
 Munson Medical Center
 Traverse City, MI
 Hemophilia Clinic of West Michigan Cancer
 Center, Kalamazoo, MI
 University of Cincinnati Medical Center
 Cincinnati, OH
 The Children's Medical Center
 Dayton, OH
 Michigan State University Comprehensive
 Center for Bleeding Disorders
 East Lansing, MI
 Children's Hospital of Michigan
 Detroit, MI
 DeVos Children's Hospital at Butterworth
 Grand Rapids, MI
 Eastern Michigan Hemophilia Treatment
 Center, Flint, MI
 Ohio State University Medical Center
 Columbus, OH

Region V-W

Mary Anne Schall, R.N., M.S.
 Northwestern University
 Chicago, IL
 Cook County Hospital - Adult
 Chicago, IL
 Children's Memorial Hospital
 Chicago, IL
 Comprehensive Bleeding Disorders Center
 Peoria, IL
 Fairview - University Medical Center
 Minneapolis, MN
 Mayo Clinic
 Rochester, MN
 MeritCare Hospital DBA Roger Maris
 Cancer Center, Fargo, ND
 Hemophilia Outreach Centre
 Green Bay, WI
 Gunderson Clinic
 LaCrosse, WI
 American Red Cross - Badger Chapter
 Madison, WI
 Rush Children's Hospital
 Chicago, IL
 Michael Reese Hospital – Adult

Chicago, IL
South Dakota Children's Specialty Clinics
Sioux Falls, SD
Comprehensive Center for Bleeding
Disorders, Milwaukee, WI
Cook County Children's Hospital
Chicago, IL

Region VI

John Drake, R.N., M.S.N.
Gulf States Hemophilia and Thrombosis
Center, Houston, TX
Louisiana Comprehensive Hemophilia
Center, New Orleans, LA
Arkansas Children's Hospital
Little Rock, AR
Oklahoma Comprehensive Hemophilia
Treatment Center, Oklahoma City, OK
Cook Children's Medical Center
Ft. Worth, TX
South Texas Comprehensive Hemophilia
Center, San Antonio, TX
Children's Medical Center
Dallas, TX
University of Texas Southwestern Medical
School, Dallas, TX

Region VII

Mike Lammer, MSW
University of Iowa Hospitals and Clinics
Iowa City, IA
Cardinal Glennon Children's Hospital
St. Louis, MO
Kansas City Regional Hemophilia Center
Kansas City, MO
Nebraska Regional Hemophilia Treatment
Center, Omaha, NE
St. Louis University Medical Center
St. Louis, MO
University of Missouri Hospital and Clinics
Columbia, MO

Region VIII

Mary Lou Damiano, R.N., M.Ed.
Mountain States Regional Hemophilia and
Thrombosis Center, Denver, CO
Ted R. Montoya Hemophilia Center
Albuquerque, NM
Mountain States Regional Hemophilia Center
Tucson, AZ
Phoenix Children's Hospital
Phoenix, AZ
Mountain States Regional Hemophilia Center
– Utah, Salt Lake City, UT

Region IX

Judith Baker, M.H.S.A.
Children's Hospital of Los Angeles
Los Angeles, CA
Alta Bates Medical Center
Berkeley, CA
University of California at Davis
Sacramento, CA
University of California, San Francisco
San Francisco, CA
Orthopaedic Hospital of Los Angeles
Los Angeles, CA
Children's Hospital, San Diego
San Diego, CA
Children's Hospital of Orange County
Orange, CA
Children's Hospital Oakland
Oakland, CA
Hemophilia and Thrombosis Center of
Nevada, Las Vegas, NV
Guam Comprehensive Hemophilia Care
Program, Agana, GU
City of Hope National Medical Center
Duarte, CA
Lucile Salter Packard Children's Hospital
at Stanford, Palo Alto, CA
University of California
San Diego, CA

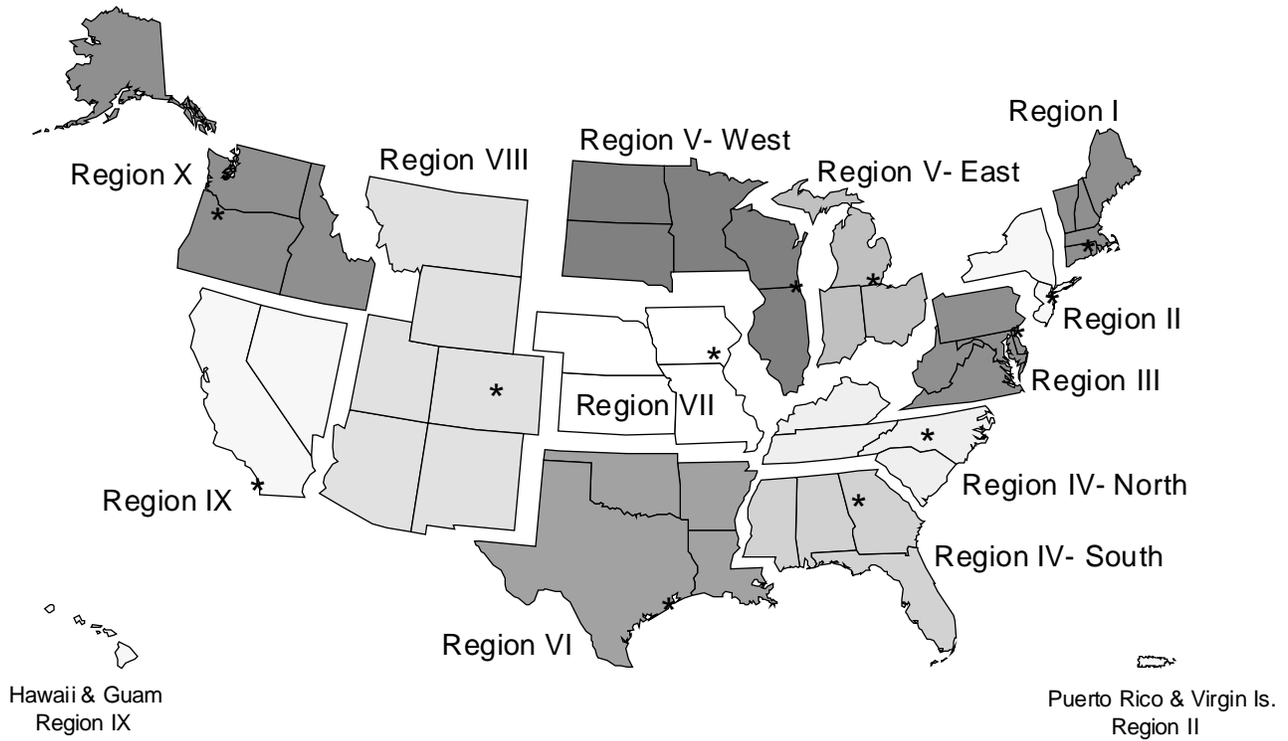
Region X

Robina Ingram-Rich, R.N., M.S., M.P.H.
Puget Sound Blood Center and Program
Seattle, WA
Oregon Hemophilia Treatment Center
Portland, OR

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Hemophilia Treatment Center Regions



*Denotes location of regional core centers.